## Peroxides. VI.<sup>2</sup> Preparation of t-Butyl Peresters and Diacyl Peroxides of Aliphatic Monobasic Acids<sup>3</sup>

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Long chain t-butyl peresters and diacyl peroxides were prepared in nearly quantitative yields and in high purity by acylation of pure t-butyl hydroperoxide and 50-65% hydrogen peroxide, respectively, in ether-pyridine solutions. These peroxides were accurately analyzed by an improved iodometric procedure employing 0.0005 and 0.002% ferric chloride hexahydrate, respectively, in glacial acetic acid. The diacyl peroxides and t-butyl peresters show an alternation in melting points.

Organic peroxides have achieved an important position in organic chemistry. Major emphasis in the past has been on the preparation and properties of aromatic and short-chain aliphatic derivatives. A detailed study of the preparation and properties of high purity long-chain peroxides, comparable to the studies on peracids,<sup>2,4</sup> was needed for clari-

(1) Eastern Utilization Research and Development Division, Agricultural Research Service, U. S. Department of Agriculture. Article not copyrighted.

(2) Paper V, This Journal, 80, 323 (1958).

(3) Presented at the Fall Meeting of the American Chemical Society, September 7-12, 1958, Chicago, Ill.

(4) (a) W. E. Parker, C. Ricciuti, C. L. Ogg and D. Swern, This Journal, 77, 4037 (1955); (b) D. Swern, L. P. Witnauer, C. R. Eddy and W. E. Parker, *ibid.*, 77, 5537 (1955).

fication of peroxide structure, systematization of their chemical and physical properties, and investigation of the kinetics of their decomposition.

Alkyl Peresters.—Milas and Surgenor<sup>5</sup> described the first synthesis and some of the properties of eight t-butyl peresters. t-Butyl perstearate was the only long-chain saturated derivative mentioned in their paper. These workers employed the Schotten-Baumann technique which has been the major preparative method in use to date.<sup>6</sup> A

(5) N. A. Milas and D. M. Surgenor, ibid., 68, 642 (1946).

(6) (a) A. T. Blomquist and I. A. Berstein, ibid., 73, 5546 (1951); (b) A. G. Davies and K. J. Hunter, J. Chem. Soc., 1808 (1953); and references contained in these papers. patent<sup>7</sup> describing a process for preparing alkyl peresters by reaction of ketenes with hydroperoxides does not cite the properties of the peresters obtainable by this process. The acylation technique in pyridine, first developed by Deninger<sup>8</sup> and amplified by Ullmann and Nadai, <sup>9</sup> was adapted for the preparation of *trans*-9-decalyl perbenzoate, <sup>10</sup> its substituted derivatives <sup>11</sup> and two esters of peroxycarbonic acids. <sup>6b, 12</sup>

The Schotten-Baumann method is inconvenient for the laboratory preparation of peresters of long-chain fatty acids because of formation of emulsions in alkaline media which are difficult to break and hydrolysis of the acyl chlorides which lowers the yield and purity of the products. In a preliminary study we acylated pure t-butyl hydroperoxide with stearoyl chloride in a solvent containing pyridine as acid acceptor and the t-butyl perstearate obtained melted 25° higher than reported. No pure long-chain t-butyl perester had apparently been prepared prior to the present investigation.

Diacyl Peroxides.—Most of the aliphatic diacyl peroxides have been mentioned in the literature, but few properties on pure derivatives have been reported. Aliphatic and aromatic peroxides, <sup>13,14</sup> fluorinated peroxides<sup>15</sup> and numerous aromatic peroxides <sup>16</sup> were obtained by acylation of hydrogen peroxide using the Schotten-Baumann method. We found the same difficulties in the application of this method to the preparation of diacyl peroxides as we observed when it was applied to the preparation of high purity long-chain t-butyl peresters.

The development of a homogeneous reaction for the preparation of diacyl peroxides requires an organic solvent for hydrogen peroxide. Ethyl ether<sup>17</sup> has been used for concentrating hydrogen peroxide from aqueous solution while a recent note by Paulsen, <sup>18</sup> in a study of the partition coefficients for aqueous hydrogen peroxide and 41 organic solvents, has shown ethers to be the best solvents. The commercial availability of high concentration hydrogen peroxide and use of diethyl ether permit direct acylation of hydrogen peroxide by a pyridine—acylation method in a homogeneous system (see Experimental.)

Acylation of peracids offers an alternate procedure for the preparation of diacyl peroxides. This method has been described earlier for preparing unsymmetrical aroyl—acyl peroxides by

(7) N. V. de Bataafsche Petroleum Maatschappij, British Patent 666,371.

(8) A. Deninger, Ber., 28, 1322 (1895).

(9) F. Ullmann and G. Nadai, ibid., 41, 1870 (1908).

(10) R. Criegee, ibid., 77, 22 (1944).

(11) (a) R. Criegee and R. Kaspar, Ann., 560, 127 (1948); (b) A. C. Cope and G. Holtzman, This Journal, 72, 3062 (1950); (c) P. D. Bartlett and J. L. Kice, ibid., 75, 5591 (1953).

(12) F. Strain, W. E. Bissinger, W. R. Dial, H. Rudoff, B. J. De-Witt, H. C. Stevens and J. H. Langston, *ibid.*, 72, 1254 (1950).

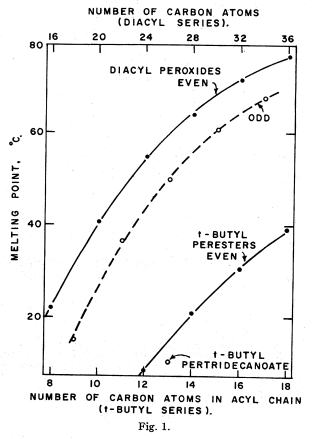
(13) W. Cooper, J. Chem. Soc., 3106 (1951).

- (14) J. Chapman and W. A. Wynne, U. S. Patent 2,771,492, Nov. 20, 1956.
  - (15) O. H. Bullitt, Jr., U. S. Patent 2,559,630, July 10, 1951.
- (16) (a) D. H. Hey and E. W. Walker, J. Chem. Soc., 2213 (1948);
  (b) A. T. Blomquist and A. J. Buselli, This Journal, 73, 3883 (1951);
  (c) F. D. Greene, ibid., 78, 2246 (1956); and references contained in these papers.
- (17) W. C. Schumb, C. N. Satterfield and R. L. Wentworth, "Hydrogen Peroxide," Monograph Series No. 128, Reinhold Publishing Corp., New York, N. Y., 1955, p. 159.

(18) F. R. Paulsen, Chemistry & Industry, 1274 (1956).

shaking a solution of the appropriate acyl chloride with perbenzoic acid in chloroform in the presence of cold alkali. 13,19 The active oxygen content of the crude products was reported to be 80–90% of theoretical. The relatively simple preparation of aliphatic peracids 4 should permit a convenient preparation of a wide variety of mixed and simple diacyl peroxides in a homogeneous medium by the pyridine modification.

Present Investigation.—Highly pure t-butyl peresters (Table I) and diacyl peroxides (Table II) of the normal aliphatic monobasic acids have been prepared in virtually quantitative yields by rapid acylation of t-butyl hydroperoxide and hydrogen peroxide, respectively, in homogeneous solution in the presence of pyridine<sup>20</sup> as the acid acceptor. The simple, rapid and accurate iodometric procedure developed for the analysis of t-butyl peresters<sup>21</sup> was adapted to the analysis of pure diacyl peroxides. A plot of melting points against carbon content (Fig. 1) shows alternation for the



diacyl peroxides and t-butyl peresters. X-Ray and polarographic data obtained for these peroxides will be reported in a subsequent publication.

Purity Determinations.—Purity of the t-butyl peresters was determined by a modified iodometric procedure in which the glacial acetic acid contained an optimum concentration of 0.002% ferric chloride hexahydrate.<sup>21</sup> Purity determinations of these

(19) H. Wieland and G. Rasuwajew, Ann., 480, 157 (1930).

(21) L. S. Silbert and D. Swern, Anal. Chem., 30, 385 (1958).

<sup>(20)</sup> On completion of this manuscript, the synthesis of several t-butyl peresters by a pyridine method was reported by P. D. Bartlett and R. R. Hiatt, This Journal, 80, 1398 (1958).

TABLE I

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Conversions and	V.HARACTERISTICS	OR LONG	L TEATNER HATTER	PERETER
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	Conver- sion, %	M.p.,* °C.	$d^{80}_{4}$	1180D	$(-\mathrm{d}n \times 10^5/\mathrm{d}t)$	Peroxide oyxgen, %21 Calcd. Found	
Perpelargonate	99	ь	0.8868	1.4271	40 (30–50°)	6.95	Found 6.93
Percaprate	95	-6.5	.88325	1.4293	39 (30–50°)	6.55	6.58
Perlaurate	99	8.0-8.6	.8784	1.4333	38 (30-50°)	5.87	5.87
Pertridecanoate		10.0-10.4	****		Anna Santa San	5.59	5.43
Permyristate	99	20.5 – 21.0	.8751	1.4368	38 (30-50°)	5.32	5.34
Perpalmitate	98	30.3-30.6		$1.4365^{c}$	38 (35-60°)	4.87	4.83
Perstearate	96	38.7-39.3°	****	$1.4386^{c}$	36 (40-60°)	4.49	4.45

<sup>&</sup>lt;sup>a</sup> Fisher-Johns melting point apparatus. <sup>b</sup> Distilled, 52-55° (10 μ). <sup>c</sup> n<sup>40</sup>D. <sup>d</sup> Reference 5 gives m.p. 14-16.5°.

TABLE II

CONVERSIONS AND CHARACTERISTICS OF LONG CHAIN DIACYL PEROXIDES

	Conver- sions.	М.р.,	Peroxide		
	%	Founda,	°C.—Reportedb	oxyge Calcd.	Found
Octanoyl <sup>d</sup>	140,445	21.8-22.4	23	5.59	5.54
Pelargonyl <sup>e</sup>	96	$13.0 - 13.5^f$	4.147.	5.09	5.05
Decanoyl	10.	40.5-41.0	المراج ورجان	4.67	4.64
Lauroyl	100	54.7 - 55.0	41 – 42	4.01	4.01
Tridecanoyl		48.3-48.8		3.75	3.73
Myristoyl	98	63.9-64.4	59	3.52	3.50
Palmitoyl	98	71.4-71.9	67-68	3.13	3.12
Stearoyl	• (•)	76.5-76.9	\$ • . • ·	2.82	2.81

\* Fisher-Johns melting point apparatus. \* Reference 13. \* Method of reference 21 but using 0.0005% FeCl<sub>3</sub>·6H<sub>2</sub>O in glacial acetic acid. \*  $d^3u_4^4$  0.9275,  $n^3v_D$  1.4363; -dn/dt 0.00027 (30–50°). \*  $d^3u_4$  0.9182:  $n^3v_D$  1.4388; -dn/dt, 0.00029 (30–50°). \* Heating curve, 13.0°, and cooling curve, 13.0–13.2°, agrees with Fisher-Johns fusion method.

derivatives by carbon and hydrogen analysis are of little value, for the presence of as much as 5% t-butyl ester will pass undetected as impurity in the corresponding perester.

Purity of the diacyl peroxides was also determined iodometrically. A brief discussion of the analysis is warranted in view of certain problems associated with iodometric methods of analysis of diacyl peroxides. Analysis of diacyl peroxides by a frequently used iodometric method<sup>22h</sup> yields peroxide oxygen values from 1–2%, and on occasion up to 5%, above calculated values. Many variations of the iodometric procedures, stated to be applicable to diacyl and diaroyl peroxides, employ modifications in solvent<sup>16b,22</sup> and in oxygen exclusion.<sup>22f,g</sup> These methods are inconvenient and lack the accuracy and precision desired for pure derivatives.

The use of ferric chloride hexahydrate in the analysis of t-butyl peresters gave such excellent results that its application to the analysis of diacyl peroxides was investigated. The results obtained on three diacyl peroxides, and one diaroyl peroxide, included to show the generality of the method, are reported in Table III. The table shows the results obtained, expressed in percentage difference from calculated, for the iodometric method,  $^{22h}$  this method modified by addition of 10 and 20 ml.

(22) (a) K. Nozaki, Ind. Eng. Chem. Anal. Ed., 18, 583 (1946); (b) C. D. Wagner, R. H. Smith and E. D. Peters, Anal. Chem., 19, 976 (1947); (c) C. G. Swain, W. H. Stockmayer and J. T. Clarke, This Journal, 73, 5426 (1950); (d) B. Barnett and W. E. Vaughan, J. Phys. Colloid Chem., 51, 942 (1947); (e) S. D. Ross and M. A. Fineman, This Journal, 73, 2176 (1951); (f) W. E. Cass, ibid., 72, 4915 (1950); (g) W. E. Cass, ibid., 68, 1976 (1946); (h) D. H. Wheeler, Oil and Soap, 9, 89 (1932).

of acetone, respectively, and the improved method developed for t-butyl peresters at varying iron chloride concentrations. The reactions were conducted in the manner reported for analysis of t-butyl peresters. <sup>21</sup> The maximum blank observed was 0.02 ml. of 0.1 N sodium thiosulfate.

## TABLE III

Comparison of Peroxide Oxygen Determined by Various Iodometric Methods

21.71.00 m	Iodo- metric	Acetone method <sup>22g</sup>		% difference from calcd. FeCl <sub>3</sub> .6H <sub>2</sub> O in acetic acid, <sup>21</sup> %		
Peroxide	method <sup>22h</sup>	10 ml.	20 ml.	0.002	0.001	$0.0005^a$
Dipalmitoyl	+2.0	+2.2	-0.10	-0.86	-1.0	-0.44
Dimyristoyl	+1.5	+0.51	+ .40	74		45
Dilauroyl	+0.50	+ .65	95	62	70	.0
Dibenzoyl	-1.0	-1.6	80	27		33
<sup>a</sup> Preferr	ed concen	tration.				

A comparison of the results in Table III clearly demonstrates that the improved method using ferric chloride in acetic acid gives results superior to the normal and acetone procedures. An optimum concentration of 0.0005% ferric chloride hexahydrate in acetic acid for diacyl peroxides, compared to an optimum concentration of 0.002% for t-butyl peresters, gives results paralleling the accuracy and precision reported for the peresters (average percentage deviation 0.29%; standard deviation 0.42%).

Additional evidence of high purity was supplied by X-ray diffraction spectra (see subsequent paper) which had sharp diffraction lines and showed no evidence of a second phase.

## Experimental

Starting Materials.—t-Butyl hydroperoxide obtained from the Lucidol Corporation, Division of Wallace and Tiernan Co., 23 was fractionally distilled in vacuo and the fractions, b.p. 42° (18 mm.), analyzing 99% or better by the iodometric procedure, 22h were retained for use. Hydrogen peroxide (50–65%) grades were generously furnished by Becco Chemical Division of Food Machinery and Chemical Corporation and E. I. du Pont de Nemours and Co. t-Butyl alcohol, Eastman Kodak White Label, was fractionally distilled, combining fractions of constant refractive index and boiling point, 82.1°. Olefin-free petroleum naphtha was obtained by treatment of the commercial petroleum naphtha, boiling range 35–60°, with concentrated sulfuric acid. The naphtha was water-washed until acid-free and distilled from anhydrous calcium chloride. Ethyl

The preparation of pure capric, lauric, myristic, palmitic and stearic acids has been described.<sup>24</sup> Pure pelargonic acid (99%) was generously supplied by Emery Industries, Inc. Acyl chlorides were prepared from the pure acids and

<sup>(23)</sup> Reference to commercial products is not intended to be a recommendation of these products by the U. S. Department of Agriculture over others not mentioned.

<sup>(24)</sup> D. Swern and E. F. Jordan, Jr., This Journal, 70, 2334 (1948).

thionyl chloride, 25 distilled in vacuo and sealed in glass am-

pules until used.

t-Butyl Peresters. Preparation of t-Butyl Perpalmitate by Schotten-Baumann Method.—To a solution of 7.3 g. (0.081 mole) of t-butyl hydroperoxide and 4.9 g. (0.27 mole) of water maintained below 20°, 20.1 g. (0.073 mole) of palmitoyl chloride and 17.7 g. of 30% aqueous potassium hydroxide solution were simultaneously added with stirring. The reaction mixture was stirred overnight, then poured into 50 ml. of ethyl ether. The emulsion was broken by neutralizing the soap and excess alkali with cold 10% hydrochloric acid solution. The ether solution was washed with three 50-ml. volumes of water, dried over anhydrous sodium sulfate, filtered and evaporated to dryness *in vacuo*. The crude product was analyzed for peroxide oxygen content by the improved iodometric method<sup>21</sup> and for acid number; peroxide oxygen, found 2.96%, calcd. 4.87%; acid number, found 89.6, equivalent to 41% palmitic acid. These analyses corresponded to 60% conversion to t-butyl perpalmitate based on palmityl chloride.

Preparation of t-Butyl Perstearate by Pyridine Acylation Method.—Stearoyl chloride (50 g., 0.165 mole) was stirred into t-butyl hydroperoxide (18.0 g., 0.20 mole) dissolved in petroleum naphtha (120 ml.) and cooled to 10°. Pyridine (15.8 g., 0.20 mole) was then added dropwise with stirring over a period of 20 minutes while the temperature was maintained below 15°. The reaction mixture was allowed to stand at room temperature for an additional 40 minutes to ensure complete reaction. The mixture was poured into a separatory funnel containing sufficient 10% hydrochloric acid solution to remove excess pyridine and pyridine hydrochloride. (Conversion and yield were determined by diluting the naphtha solution to a known volume, pipetting a sample for iodometric analysis and drying the sample under vacuum to remove solvent and unreacted hydroperoxide.) The petroleum naphtha solution was washed with 5% potassium bicarbonate solution, water-washed, and dried over anhydrous sodium sulfate. After filtration, t-butyl per-stearate crystallized at 2° in long, hard needles (Table I).

Other t-Butyl Peresters.—The other t-butyl peresters listed in Table I were prepared by the pyridine acylation procedure described for t-butyl perstearate, with the quantities of reactants as shown (recrystallization temperature is given in parentheses): t-butyl perpelargonate: 44.3 g. (0.25 mole) of pelargonyl chloride, 27.0 g. (0.30 mole) of t-butyl hydroperoxide, 23.7 g. (0.30 mole) of pyridine (perester distilled from alembic apparatus, head temperature 55–58°/10 nydroperoxide, 23.7 g. (0.30 mole) of pyridine (perester distilled from alembic apparatus, head temperature 55–58°/10 μ); t-butyl percaprate: 50 g. (0.262 mole) of capryl chloride, 28.3 g. (0.314 mole) of t-butyl hydroperoxide, 24.8 g. (0.314 mole) of pyridine (−40°); t-butyl perlaurate: 34.2 g. (0.156 mole) of lauroyl chloride, 17.0 g. (0.188 mole) of t-butyl hydroperoxide, 14.9 g. (0.188 mole) of pyridine (−35°); t-butyl tridecanoate: 2.6 g. (0.011 mole) of tridecanoyl chloride, 1.6 g. (0.018 mole) of t-butyl hydroperoxide, 1.2 g. (0.015 mole) of pyridine (−16°); t-butyl permyristate: 50 g. (0.203 mole) of myristoyl chloride, 21.9 g. (0.243 mole) of t-butyl hydroperoxide, 19.2 g. (0.243 mole) of pyridine (−20°); t-butyl perpalmitate: 41.2 g. (0.15 mole) of palmitoyl chloride, 20.3 g. (0.225 mole) of t-butyl hydroperoxide, 15.0 g. (0.188 mole) of pyridine (2°).

Diacyl Peroxides. Preparation of Dipalmitoyl Peroxide by Schotten-Baumann Method.—Palmitoyl chloride (20 g., 0.073 mole) was dissolved in chloroform (20 ml.) and cooled to 5°. Hydrogen peroxide (2.5 g. of 50% concentration, 0.037 mole) and sodium hydroxide (11.6 g. of 25% concentration) were simultaneously added dropwise with

concentration) were simultaneously added dropwise with vigorous stirring, while the temperature was maintained below 15°. The mixture was stirred for one nour area complete addition of reactants and poured into a chlororeactants and poured into a chioroform-water mixture. The emulsion could not be broken without partial acidification. The chloroform solution was washed with water, dried over anhydrous sodium sulfate, filtered and evaporated. The product, recovered in 77% yield, had a peroxide oxygen content 64% of calculated. This represents a 50% conversion based on palmitoyl chloride. The resulting product was difficult to purify by direct crystallization. crystallization.

Preparation of Dimyristoyl Peroxide by Pyridine Acylation Method.-Only the preferred procedure will be described in detail: myristoyl chloride (24.7 g., 0.10 mole) was dissolved in ether (175 ml.) and cooled to  $0^{\circ}$ . Hydrogen peroxide (4.25 g. of 60% concentration, 0.075 mole) was added, followed by the dropwise addition of pyridine (9.5 g., 0.12 mole) while the temperature was maintained at 0-5°. The diacyl peroxide precipitated as it formed. After complete addition of pyridine, the ice-bath was removed and stirring of the slurry was continued for a total time of one hour. A homogeneous solution was obtained at room temperature by addition of more ether. The ether solution was washed with dilute hydrochloric acid, 5% potassium bicarbonate, followed by water and then dried over anhydrous sodium sulfate. The solution was diluted to a known volume and a sample pipetted for iodometric analy-This procedure was followed for a series of experiments varying the amount of solvent, temperature of reaction, mole ratio of reactants and time as summarized in Table IV. For product isolation, the original ether solution was cooled to 0-2°.

The temperature at which reaction is conducted is the most important factor for attainment of high conversions. Sufficient solvent is used to permit unhindered stirring of the slurry which forms by crystallization of diacyl peroxide during reaction. The reaction is believed to be instantaneous, but an additional time is allowed after complete pyridine addition to permit any acyl chloride occluded by the precipitating peroxide to redissolve and effect quantitative reaction.

## TABLE IV

DETERMINATION OF CONDITIONS FOR MAXIMUM CONVER-SION OF MYRISTOYL CHLORIDE TO DIMYRISTOYL PEROXIDE Mole ratio

of reactants, RCOCl <sup>a</sup> : pyridine: H <sub>2</sub> O <sub>2</sub>	Solvent, ml.	Temp.,	Time,	Conversion,
1:1.1:0.55	CHCl <sub>3</sub> , 75	15-20	3	65
1:1.25:0.55	CHCl <sub>3</sub> -ether 1:2, 112	15-20	1	71
1:1.3:0.75	CHCl <sub>3</sub> -ether 1:14,			
	188	15-20	2	90
1:1.1:0.60	Ether, 175	0-5	0.5	84
1:1.2:0.75	Ether, 175	0-5	1	98

<sup>a</sup> 0.10 mole.

Other Diacyl Peroxides.—Chloroform (75 ml.) was used as a cosolvent at the end of the reaction period only for the relatively insoluble dipalmitoyl and distearoyl peroxides. Distearoyl and dipalmitoyl peroxide were crystallized from the chloroform-ether solution. Ditridecanoyl and dilauroyl peroxides were crystallized from ether. Didecanoyl and dipelargonyl peroxides were crystallized from olefin-free petroleum ether.

Acviation of Perlauric Acid.—Perlauric acid (4.80 g., 0.0222 mole) was dissolved in ethyl ether (54 ml.), cooled to 0° and lauroyl chloride (4.87 g., 0.0222 mole) was added. Pyridine (2.1 g., 0.0266 mole) was added dropwise while stirring the mixture at 0-5°. The mixture was stirred for an additional 40 minutes while it was allowed to warm to room temperature. The ether solution was washed with water, dried over anhydrous sodium sulfate, and analyzed iodometrically. Peroxide oxygen was 89.3% of calculated.

The above experiment was repeated at 10-15°. At the higher reaction temperature, the peroxide oxygen content of the product was only 79.4% of calculated. Recrystallization of both crude products from ethyl ether-petroleum naphtha (1:1) yielded pure dilauroyl peroxide.

Melting Points.—Melting points were determined on a Fisher-Johns melting point apparatus. Crystals with melting points between ice and room temperatures were determined on the Fisher-Johns block in a cold room thermostatically controlled at 2°. The compounds melted sharply with controlled heating of the block. The accuracy of the method was checked by comparing the value found for dipelargonyl peroxide crystals with the value determined from its heating and cooling curves using a calibrated N.B.S. thermometer.